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APPLICATION NUMBER: 60/391,575

FILING DATE: June 27, 2002

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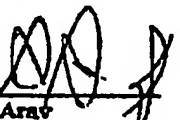
EMBRYO GUARD

described and claimed in the following patent applications:

U.S. Provisional Application identified as Attorney docket No. 791/14 and executed the same date as this assignment;

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Signed and sealed this 22 day of Temp 2002


Amir Arav
19/10/2002


Meir Uri

60391575 .06

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As-Filed New Application

Level 1
Version 1.1
Updated - 8/01/01

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Transmittal



Level - 2
Version 1.1
Updated - 8/01/01

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Docto Number: 741114

PROVISIONAL APPLICATION FOR PATENT COVER SHEET

This is a request for filing a PROVISIONAL APPLICATION FOR PATENT under 37 CFR 1.53 (c).

INVENTOR(S)/APPLICANT(S)

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Additional inventors are being named on Page 2 attached hereto

Title of the Invention (Up to characters max)

EMBRYO GUARD

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Country _____

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ENCLOSED APPLICATION PARTS (check all that apply)

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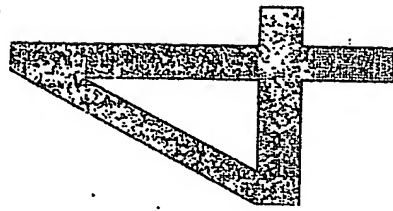
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(Page 1 of 1)

J1129 U.S. PTO
06/27/02



Specification



Level 2
Version 1.1
Updated - 8/01/01



June 2002

IMT Ltd.

Microscopic Monitoring

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Technology and Products

Microscopic Monitoring

During its many years of research in the fields of cryopreservation and reproduction, IMT has developed several techniques and supporting technologies to improve its research process. The MMS technology is a result of one of these developments. Initially, the company's researchers needed a tool that allows microscopic monitoring of cells during the freezing process. The solution was a miniature device, consisting of a CCD camera, a microscopic long-distance objective lens and a special adapter. That was the beginning of a new field for IMT and based on that simple device, the company develops several important products, including the EmbryoGuard.

1. The Technology

Microscopic monitoring is very common in biology, as it is a basic tool for most procedures. In every lab there are usually several types of microscopes and most of the devices have a special adapter for video or CCD camera. However, this simple approach has several limitations. First, the image must go through the optical system of the microscope. This means that the microscope should be a high quality device, so that the quality of imaging remains untouched. Such high quality microscopes are very costly. In addition, the size of the microscope limits its applications. If one wants to monitor cell development *inside* an incubator, he can "build" an incubator that surrounds an existing microscope, or he can take the cells out of the incubator and place them under a microscope in a warm environment. IMT has succeeded in miniaturizing the whole system, while avoiding the need for a microscope. A very small CCD or video camera, with a special adapter and a microscopic lens, can provide the same results as an expensive and high quality system.

An improvement to this basic technology was introduced when the Company developed a robotic system that can control multi-sample monitoring, with X-Y-Z micro movement. This system is based on one or two microscopic CCD cameras, together with robotic features that can move the cameras and the samples. This system is then placed *inside* a standard incubator. A control unit placed *outside* the incubator helps to control focusing and illumination, changing between samples, and more. The

control unit can be handled manually or by computer software and the images are screened on a standard monitor.

2. Products and Applications

The Company is developing several products based on that technology. The main one is the Embryo Guard.

The Embryo Guard (EG) is a robotic system for microscopic monitoring and control over embryo development during IVF procedure. It has computerized control and software that assists with embryo evaluation, supports the selection process, and controls the matching process (patient/ovary/sperm/embryo).

During IVF procedure, it is extremely important to monitor the development of fertilized eggs, from the moment of fertilization up to the stage when 2-3 embryos are selected for transplantation. The importance of this monitoring is derived from the fact that eventually, the best embryos should be chosen for transplantation. Therefore, the clinician must watch very carefully every development stage of the embryos.

Problems being addressed

Change in conditions - Currently, the monitoring procedure is done manually, by taking the embryos out of the incubator, placing them under a microscope and investigating their development.

This approach has several disadvantages that usually damage the embryos. To assure the best conditions for embryo development, it is essential that the embryo remains in a stable controlled environment, as provided by the incubator. Any change in these conditions can easily harm the embryo. Therefore, the procedure of taking the embryo out of the incubator, although it is necessary, has a bad effect on the embryo development. In addition, the optimal way to evaluate embryo development is to monitor it every 3 hours, but again, since this monitoring might be too risky, most IVF labs prefer to perform this evaluation much less frequently. Another problem raised from the need to monitor an embryo under a microscope is that the embryo must be under a special solution (oil) that can damage it.

The EG solves these problems by providing continuous monitoring of the embryo, without taking it out of the incubator. The EG, automatically, monitors each embryo

every 3 hours, or continuously (time-lapse recording) and stores this data (as image files) on the embryo records. This process is done inside the incubator which means that embryos do not experience any change in condition and also, there is no need to use oil or any other solution that may cause damage.

Another important advantage of the EG is the ability to control it from a distance, using the Internet. The embryo specialist does not have to be present in the IVF lab, and he can control the whole process using a standard computer connected to the Internet.

Standards and data recording - Another problem addressed is the lack of standards and data recording. Currently, there is no software application that supports the IVF procedure in terms of embryo development and selection of eggs and embryo. Without such supporting software, IVF labs collect data on the embryos in a variety of ways, with no specific standard or quality control, and in addition, most of this data collection is done by paper work.

The EG includes a software that satisfies these needs. Each embryo in the incubator has its own record, containing all the information from the initial stage. The software also automatically collects and stores pictures of each embryo in each stage. In addition to data collecting, the software also helps to evaluate the embryo by indicating in which stage it should be, how many cells it should have, what should be the next stage and the timing for this stage, etc. It can also provide a multi-embryo screen that helps to compare their visual shape.

Matching - IVF labs pay a lot of attention to the issue of matching between oocytes and sperm, or embryos and patient. Even a minor mistake could be a personal disaster for the future parents and a major legal problem to the lab. One component of the EG aims to solve this problem. The EG has a unique matching system that makes sure that no mistake can happen. Every sperm sample and every oocyte are labeled (on the dish) with a barcode labeling system. This procedure is then stored in the computer record as the first step of the IVF procedure. From the moment of labeling, every procedure must first go through a barcode reader that stores the information under the patient record. Before fertilization, the EG software identifies both patients and indicates if there is matching or not. The fertilized eggs are then placed inside the incubator (again - after going through the barcode reader). If during the incubation

period, or even before transplanting the embryos, the clinician needs to take the embryos outside the incubator, the software identifies the specific dish and let the clinician take only this one.

Schedule

IMT plan to introduce the first commercial version of the EG on July 2002.

Consumable products

The Embryo Guard can handle up to 12(?) dishes simultaneously. Each dish is for one specific patient and it can contain up to 10(?) embryos. The dish is a regular and standard dish, sterilized, and is currently available in the market. For locating and identifying the embryos in specific locations around the dish, the company offers a special sticker for each dish, which also contains barcode ID. The EG cannot operate without IMT's stickers.

embryo guard notes

Embryo Guard

Notes from interview with Amir Arav, 26 June 27-06-2002

Based on US 6,166,761. Now full robotic system.

Point 1: Follow development of embryos because in vitro fertilization depends on first cleavage. Need to know timing of cleavage. Implant three-day-old embryos. Hard to tell which one to pick.

Point 2:

Prior art: embryos removed from incubator to exchange medium.
New: automatic medium exchange. Add medium, remove medium, or do both. Gass the medium before or after warming up the medium.

Point 3: Zona pellucida stays thick in an incubator. As a result, the embryo may die. Focus a laser beam through a microscope to heat the zona pellucida to cut it open. Do this inside the incubator. Cutting may be done manually by technician, or automatically. Prior art is to remove embryo from incubator and cut zona pellucida outside incubator.

Point 4: Fluorescent markers. Also for preimplantation diagnostics.

Point 5: Insemination inside incubator.

Point 6: Preparation for cryopreservation: is a special case of Point 1.

In general: micromanipulation of oocytes and embryos is done inside the incubator.

Matching using bar code (or equivalent: remotely readable chip, imaging, etc.)

Identify gametes, oocytes, sperm upon collection.
Stickers on test tubes, vials, petri dishes (containers generally). ID text (parents names etc.,) matched automatically to ID code.

Box outside incubator has place for one test tube and one petri dish.
Match test tube to petri dish based on bar code. Match embryos and cryogenic vials (need liquid-nitrogen-resistant bar code) for cryopreservation. Match again when transferring embryos to womb.
Gamete Intra Fallopian Transfer. Zygote Intra Fallopian Transfer. Match at Pre-Implantation Diagnosis.

Management software: tells you what to do when (timing is critical). Warns if embryos are outside incubator too long. Collect history automatically.

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Patent on EmbryoGuard

1. On line monitoring, live lens recording, medium change over and assisted hatching of embryos inside the incubator.

It is well recognized that the timing of the first cleavage and the morphology of the embryos determine the successful of the IVF procedure. In addition, change over of medium and assisted hatching are other reasons for removal the embryos from the incubator which could be optimized if they could be performed inside the incubator. Many opening of the incubator i.e. for microscopic evaluation, medium change-over and assisted hatching, will affect the embryo condition (temperature, gas concentration and humidity).

We describe here a robotic system which will operate in the incubator with the following feature:

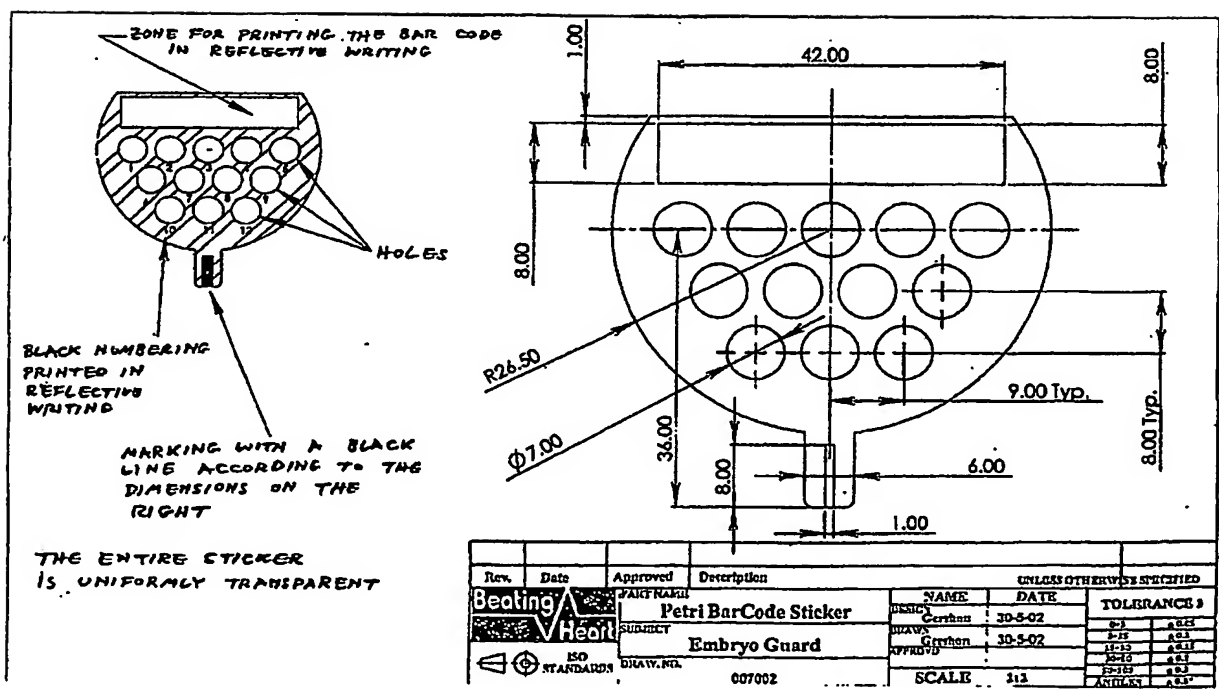
1. A microscopic follow up of the development of embryos inside the incubator with the following possibility. A. A real-time evaluation of the embryos using up to 4 different microscope CCDs which could be operate on 3D movements also by using internet compatibility. B. automatic photographing system for image of up to 12 different dishes in which 12 drops are placed in each of the Petri dish.
2. Medium change over is done by computerized injection to each of the drops with small volume of 1-10 microliter of fresh medium which is maintained cooled before injection and then warmed up, gassed and added to the drop.
3. Assisted hatching is done by laser beam performing on the zona pelucida in order to assist blastocyst hatching.
4. Evaluation using fluorescent markers which are loaded with the injectors and are detected by the embryoguard.
5. Insemination could be done stepwise by the injectors insert sperms directly to the drops.
6. Preparation oocytes or embryos for cryopreservation inside the drops using the computerized injector in a stepwise manner and according to the centric behavior of the cells.

2. Control matching using barcode system.

It is estimate that there are hundreds of mistakes in IVF matching worldwide. Identification of embryos is done by the technicians created humanity mistakes. We propose of using a adhesive sticker with a barcode for test tube and Petri dish. The EmbryoGuard read the barcode and identify the oocyte for matching with sperm. In a separate apparatus which is placed outside of the incubator (en EmbryoSealer). The matching could be performed in several levels:

1. inside the incubator before the oocytes are fertilized.
2. when sperm arrive to the lab
3. between sperm and oocytes
4. for PGD
5. for cryopreservation

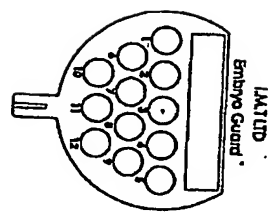
In a case of no matching the EmbryoGuard will not aloud to be open and remove out the oocytes or embryos or done any other function.



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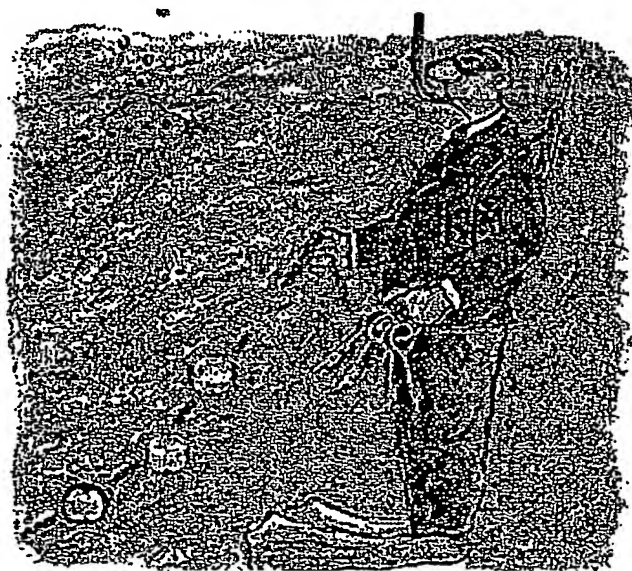
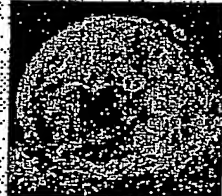
DR. M. FRIEDMAN +++ BILL

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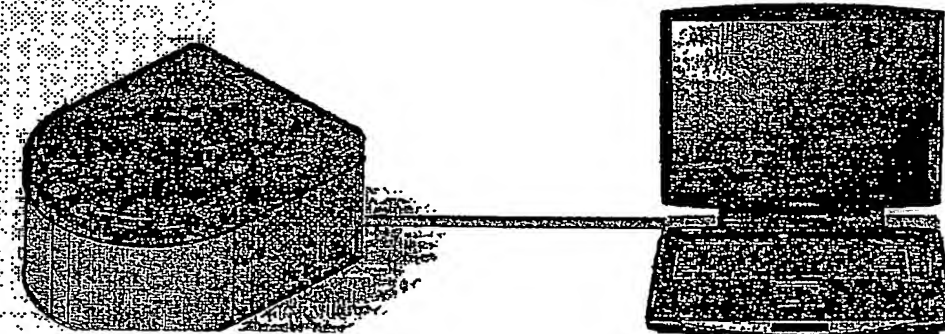
The EMBRYOGUARD



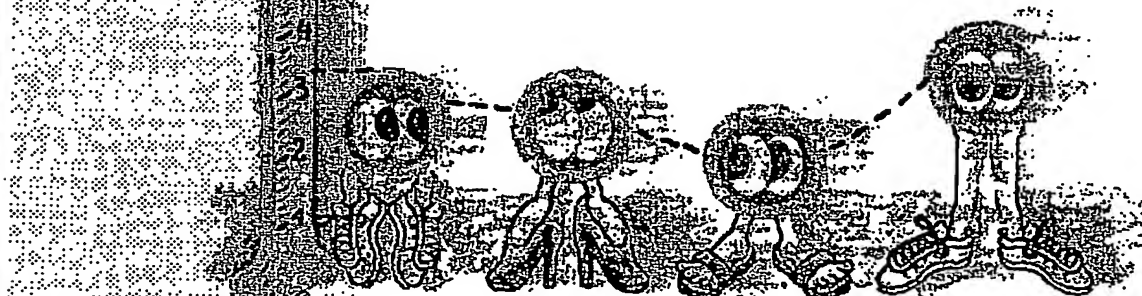
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The Embryoguard

- A new system which contains three microscopic CCD cameras located inside the IVF incubator, including active matching management tool

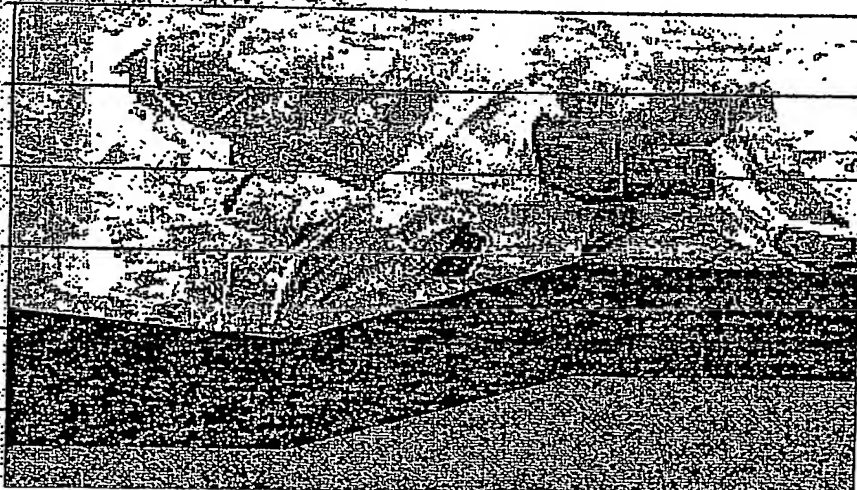


On-line monitoring & time-laps evaluation of embryos inside incubator



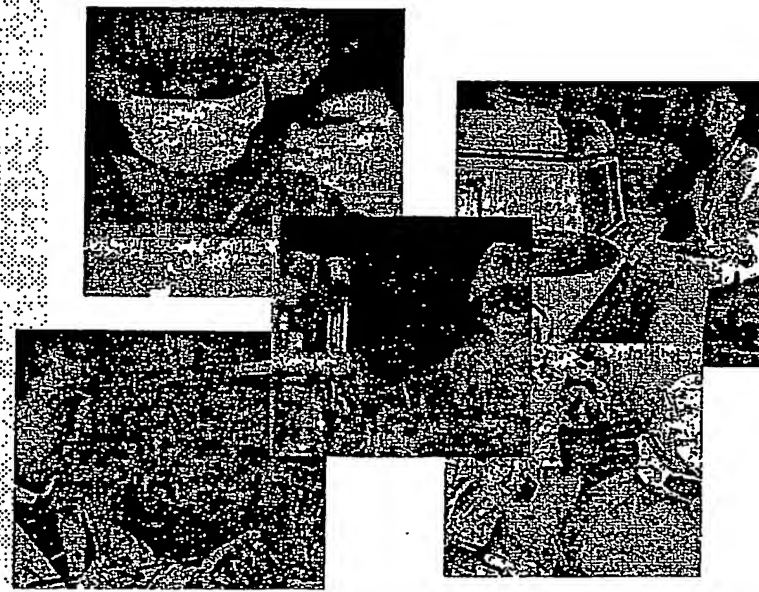
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**Improved success rates by selecting of
embryos based on cleavage timing**

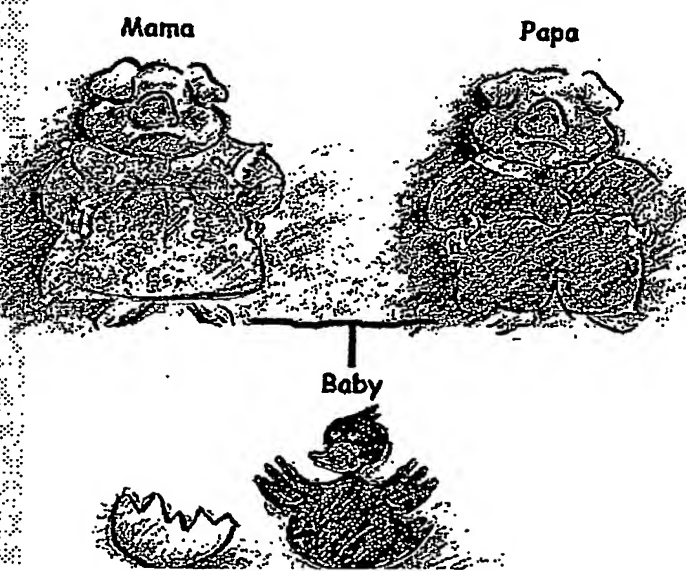


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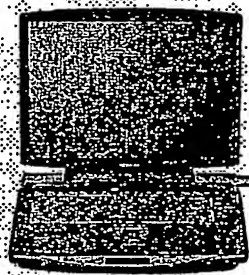
Optimal management of IVF lab procedures



Controlled matching utilizing the Barcode identification system

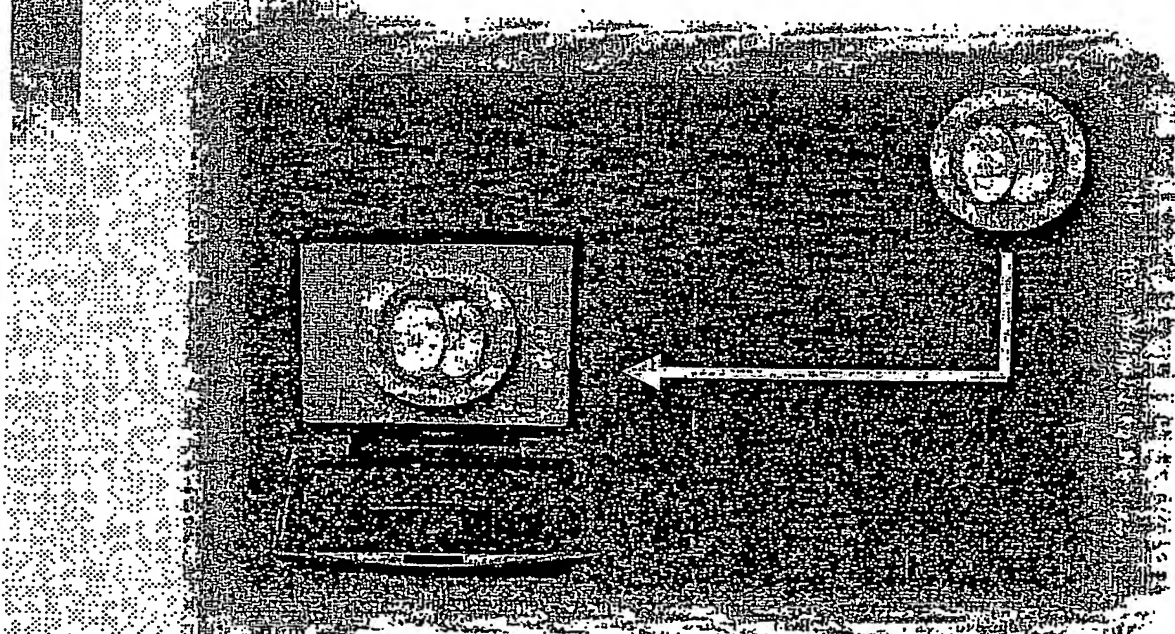


Complete documentation



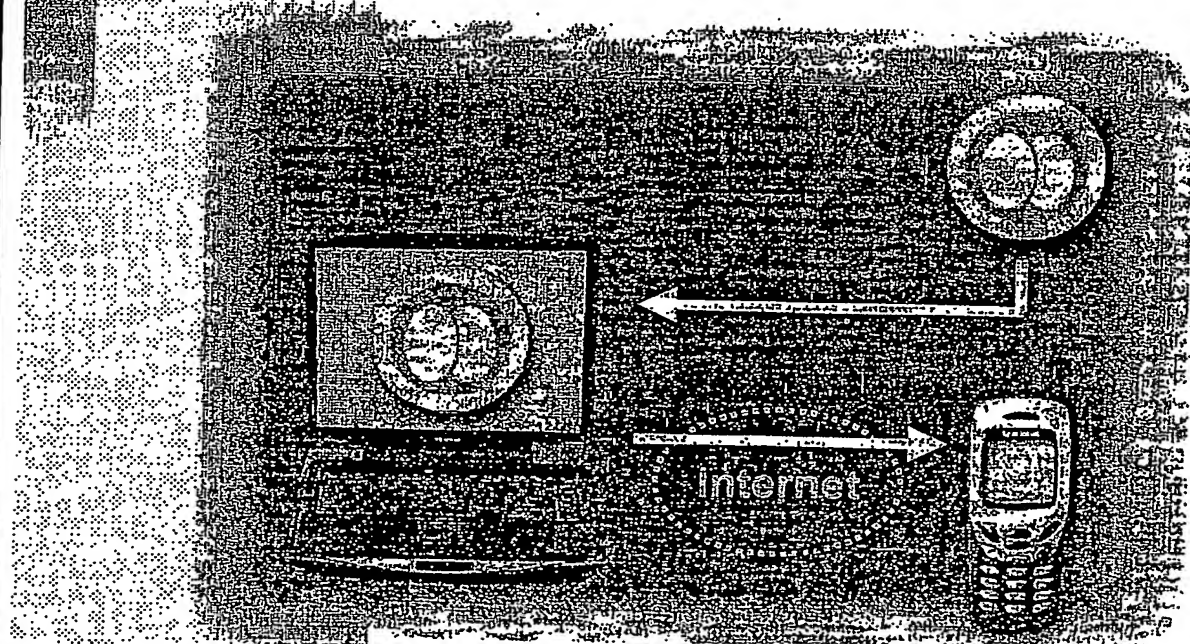
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On line control per procedure



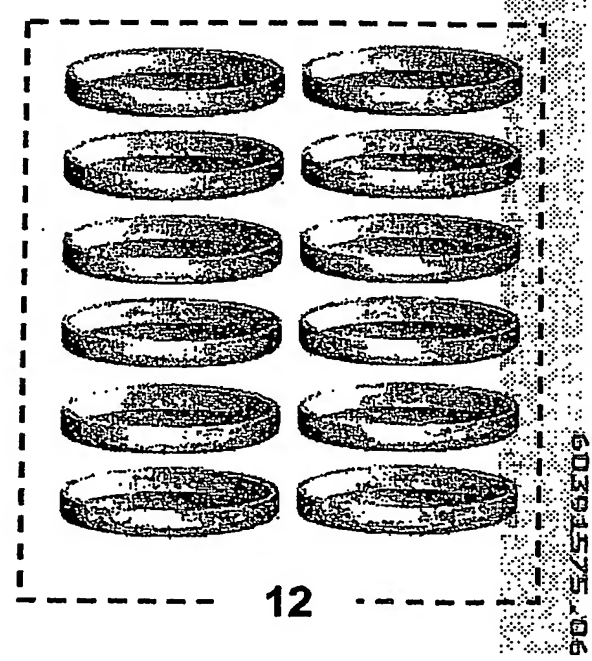
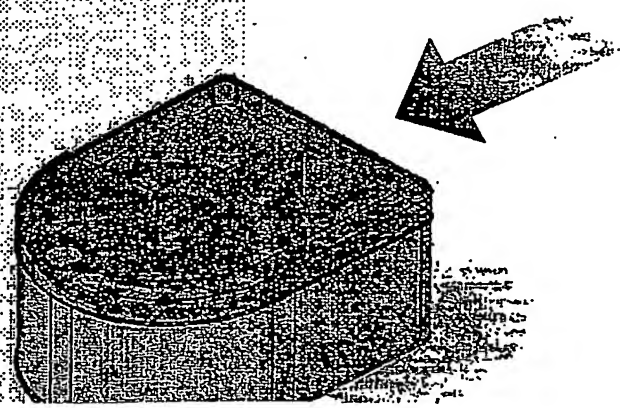
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Internet compatibility

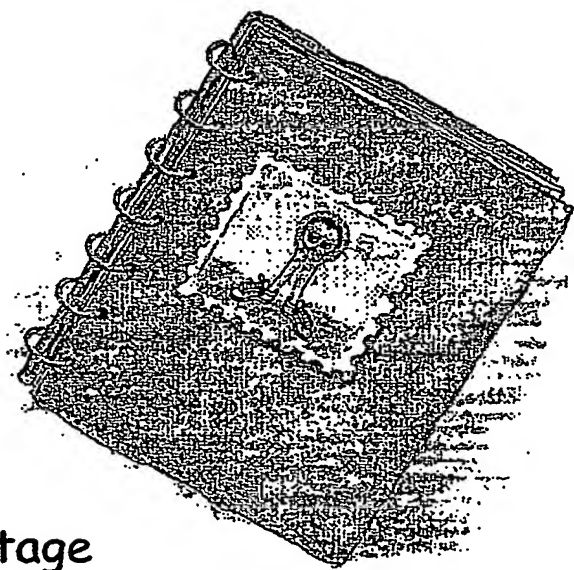


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Real time evaluation of up to 12 dishes simultaneously

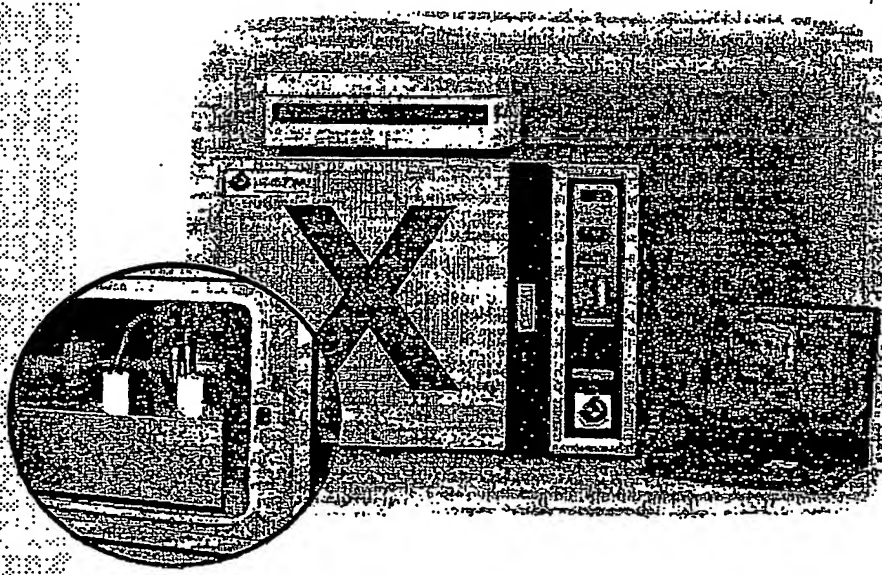


**Full robotic 3D movement of CCD
microscopic cameras inside the incubator**



**Start your
photo album
from your 2PN stage**

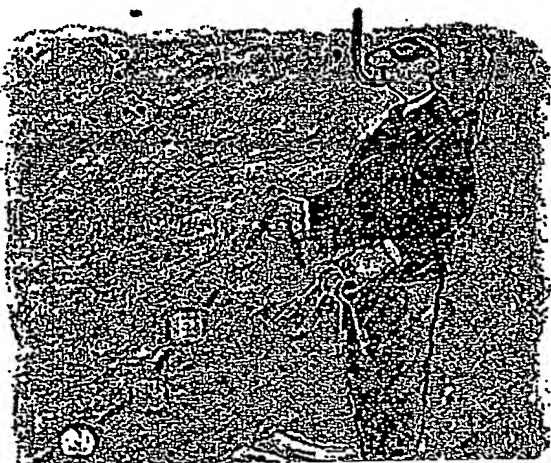
Reduces the need opening the incubator

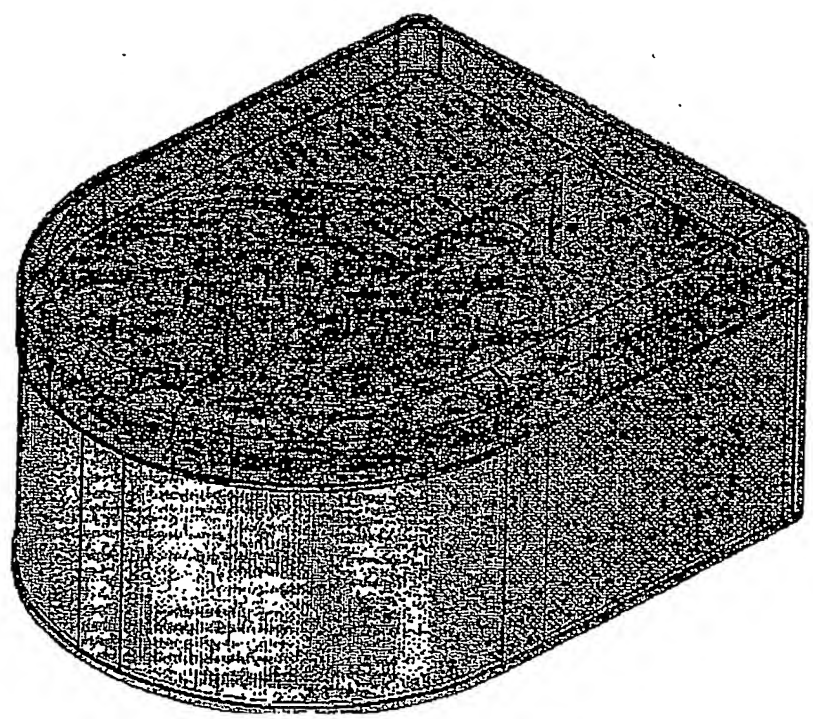


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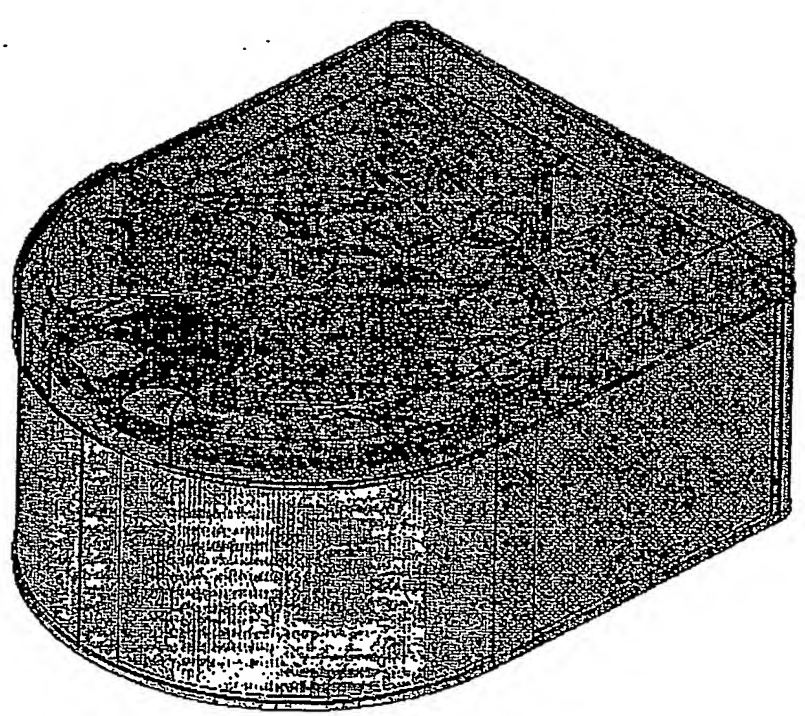
For More Information

- E-mail: embryoguard@cryo-imt.com
- Site: www.cryo-imt.com

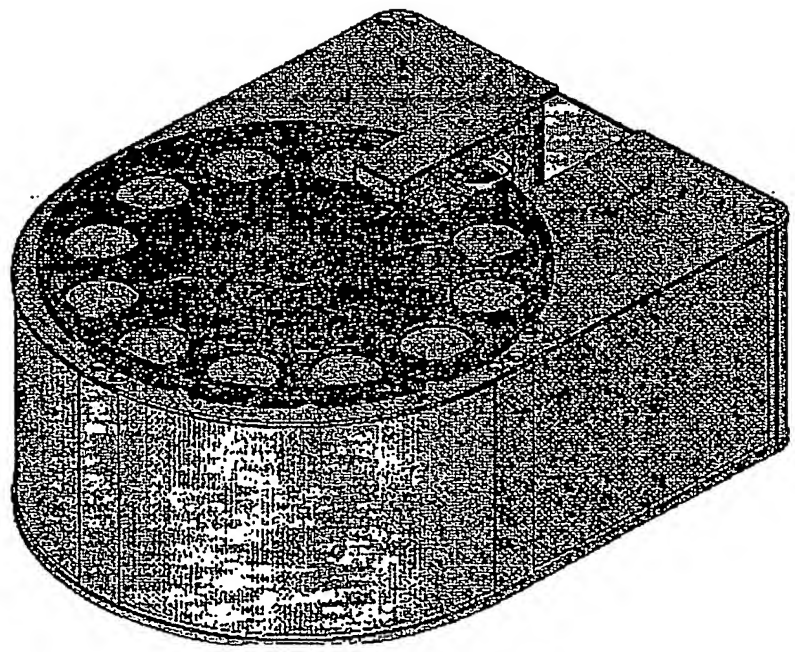




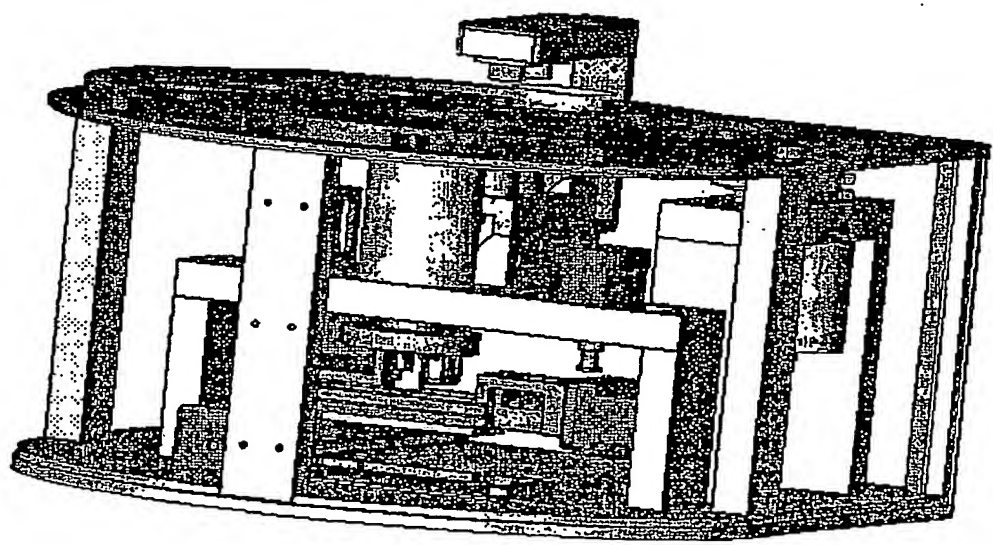
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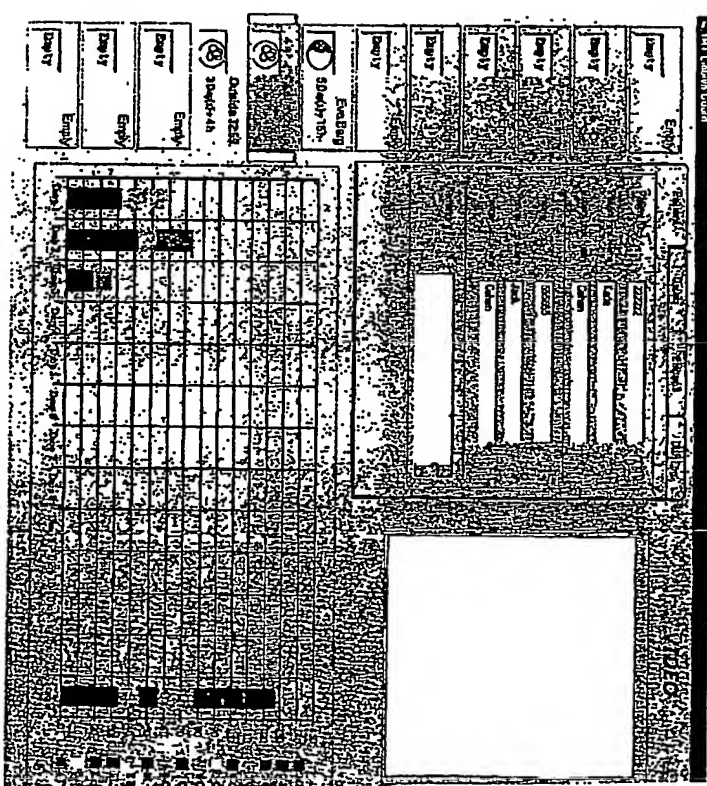
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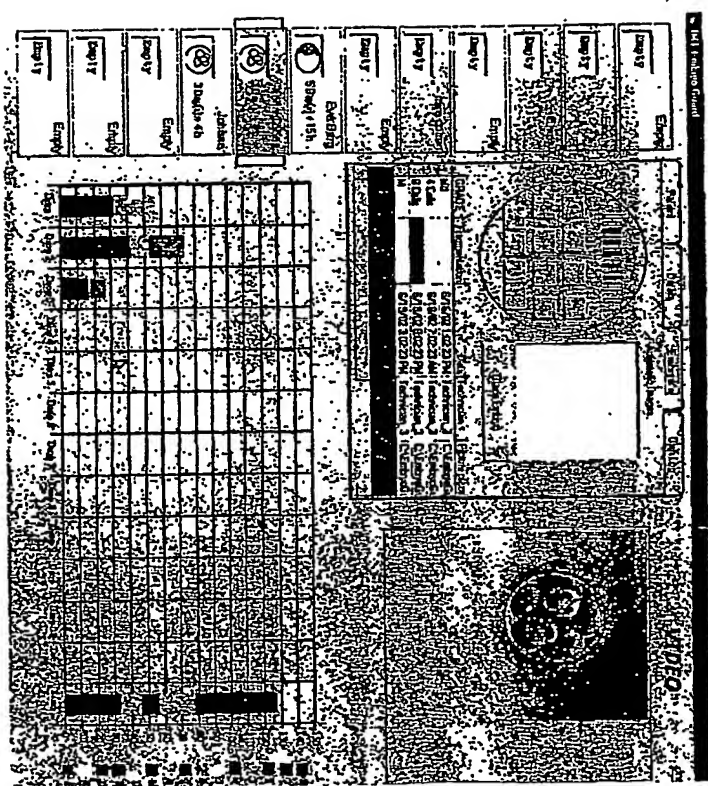


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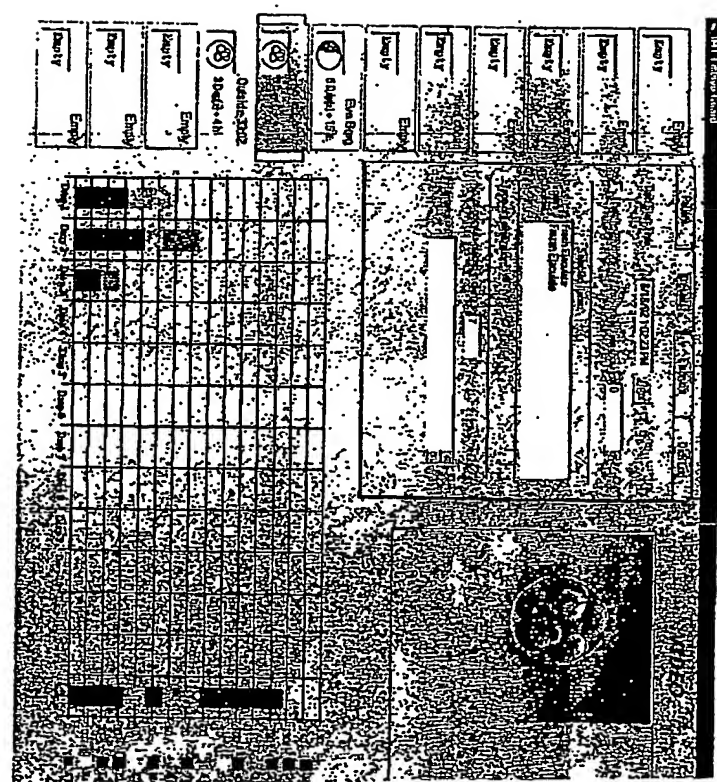


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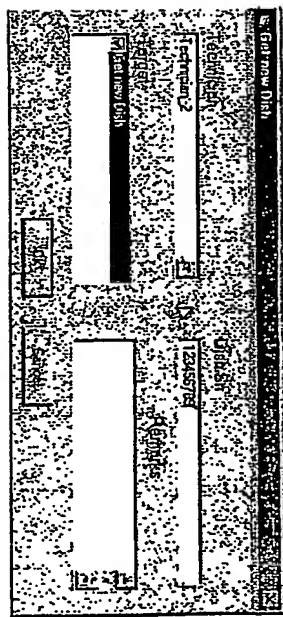




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2. Patient Name: mika; Serial Num: 2333333

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5.1.1. *Bovine Semen*

The worldwide use of bovine semen in animal husbandry is enormous, with approximately 100 million doses administered each year. With costs per dose ranging between approximately \$4-50/dose, and with a mean of about \$10/dose, this total industry has a value of about \$ 1 billion per annum. IMT estimates that at a charge of \$0.90 per dose, its market share would be about \$100 million per annum.

Use of IMT's MTG™ 525 equipment in the freezing of bull semen has been shown to yield a greater proportion of viable sperm than other conventional methods. This leads to more doses of semen available per ejaculate, a major commercial advantage for the user. In addition, because the IMT technology reduces significantly the damage to sperm cells, it may be possible to achieve successful insemination with smaller doses of semen, an additional advantage of this technology. Field trials are now underway at Cogent Breeding Ltd (UK) to demonstrate this added advantage. Additional potential advantages of IMT technology - the ability to freeze larger volumes of semen in a single container, the ability to re-freeze semen specimens after initial thawing, and the freezing of semen specimens that have been sex-sorted - are some additional applications that are being investigated by IMT. All these variations are desirable additional applications of the IMT technology that would have significant operational and commercial benefit to the user. By developing and demonstrating these applications, IMT will be able to secure its place in the market for a long time.

IMT currently has the MTG™ 525 module in routine use in the UK (Cogent). It is also under evaluation in Switzerland. IMT anticipates that an additional 5-6 machines will be placed during the second half of the year 2002.

5.1.2. *Equine Semen*

Equine semen freezing is not as far advanced in the world market as is bovine semen, but the worldwide market is growing rapidly. Stallion semen is more difficult to freeze than bovine semen, and the semen of about 25% of stallions seem to be resistant to freezing. Success rates for equine artificial insemination is about 40-50%.

As currently practiced, stallions selected for breeding are often shipped at sometimes great distances so that natural insemination can take place. This approach is costly and inefficient. In contrast, artificial insemination permits the "banking" of selected stallion semen. Insemination of a mare can take place at the convenience of the owner, even long after the stallion has died. Thus, the use of semen storage offers the horse's owner a form of insurance over his (usually large) investment.

Preliminary studies indicate that IMT freezing technology can improve semen recovery by about 20%. More importantly, semen from stallions that could not be frozen previously has been successfully frozen by this technology and has resulted in pregnancies. This opens up a large and remunerative field for IMT's technology. Based on a \$20/dose charge, IMT estimates this current annual market at about \$100 million. IMT, in collaboration with Cogent is now continuing to perform field trials in Europe to further document the efficacy of its equipment in the freezing of equine semen. Results are anticipated by the end of year 2002.

5.1.3 *Porcine Semen*

IMT has recently added the freezing of porcine semen to its growing list of animal artificial insemination applications. Studies in IMT's laboratories indicate that, when using the MTC™ 125 module, there is an excellent, 95% recovery of viable sperm. Field trials in which porcine semen frozen with IMT's technology will be used to inseminate and produce viable offspring are scheduled to commence in the Summer of 2002.

Average production cost of one insemination dose is about \$3.5 when produce in the farm (disregarding the boar's genetics). An average farm produces about 15,000 doses annually. This is a market that is in its relatively early stages of development and its size is difficult to estimate. IMT anticipates that an MTC™ technology will change this field and that there will be newly established artificial insemination farms supplying frozen porcine semen. IMT estimates that it will be able to charge a royalty fee of \$0.90 per dose of porcine semen frozen.



Successful pregnancies in cows following double freezing of a large volume of semen

Abstract

The objective of the following paper is to describe a new technology for large volume and double freezing of semen in 12 ml test tube. Semen from two different bulls was frozen with a new technique using 12 ml test tube and was refrozen after thawing in mini straws. All freezing was done in a "Multi thermal gradient" (MTG) freezing apparatus, which moves the container at a constant velocity (V) through a thermal gradient (G) producing a controlled cooling rate $B = (G) \times (V)$.

Each of the two bulls ejaculated were evaluated for post thaw motility in the lab and then in a field trial which was carried out in a split sample mode. We inseminated 105 cows after double freezing/thawing cycle, and another 123 cows were inseminated with semen frozen in mini-straws and a conventional method. Results showed a 75±5% post thaw motility after freezing a 12ml test tube and 50±5% after second freezing/thawing in mini-straws, respectively. Controlled vapour freezing showed a 60±10% post thaw motility. Results of the field trial showed a pregnancy rate of 44% (47/105) for the double freezing group in comparison to 45.5% (56/123) for the controlled group. These results can be beneficial for large volume freezing, and therefore for bull semen cryobanking in a large volume which will be followed by second freezing in a regular insemination volume.

Bull semen Cryobanking

Cryobanking of semen has had a major impact on dairy cattle genetic breeding. In addition to its role in young bull genetic breeding, cryobanking of bull semen is an important backup for sufficient insemination doses in cases of disease, infertility or mortality.

Freezing and storage of semen is done regularly using mini (1cc) or midi (5cc) straws. However, cryobanking of a large number of straws is time consuming, expensive and requires a lot of storage space and liquid nitrogen. An alternative procedure which will reduce these expenses could be the freezing of a whole ejaculate in one test tube (12ml) and, only when needed (when the bull is a "proven bull") the test tube will be thawed and then refrozen in regular mini straws. We describe here the use of a new technology for large volume (whole ejaculate) freezing/thawing and refreezing in mini-straws.

MTG technology

Our novel freezing technology is based on "Multi-thermal gradient (MTG®) IMT, Israel) (1) directional solidification and is used mainly for freezing sperm and large

tissue. The semen in the test tube is moved at a constant velocity (V) through a linear temperature gradient (G), so the cooling rate ($G \times V$) and ice front propagation are precisely controlled (Fig. 1).

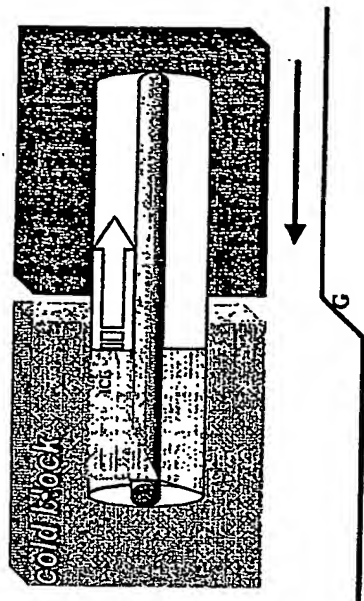


Fig. 1 Schematic design of the MTG freezing

This method also enables the incorporation of controlled seeding into the freezing process. When any liquid is cooled below its freezing point, it remains a liquid, in an unstable super-cooled state, until freezing starts at randomly distributed nucleation sites and spreads throughout the entire volume of the liquid. As discussed above, in the conventional equilibrated method of freezing, ice grows with uncontrolled velocity and morphology, and may disrupt and kill the cells of the samples. Ideally, the velocity of the freezing front should be such that the ice morphology does not disrupt the cells or tissue. However, the rate of cooling appropriate for favorable ice morphology may not be appropriate for other desired outcomes of a sample's freezing protocol. The laterally varying gradient used in our technology allows cooling to proceed at differing rates under varied temperature regimes, thereby facilitating full control over nucleation and ice crystal morphology. This technique allows very precise control of the cooling rate (0.01 to 100°C/minute) within a large volume.

The freezing apparatus can control ice crystal propagation by changing the thermal gradient (G) or the liquid-ice interface velocity (V) and so optimizing the ice crystal morphology during freezing of cells and tissue. The rate of cooling also affects the

morphology of the intercellular ice crystals (3): morphologies such as closely packed needles kill cells by external mechanical damage (unpublished observation). Thus, maximizing the survival ratio of cells subjected to freezing and thawing requires careful control of the freezing process, i.e. interface velocity. Using a cryomicroscopy observation we found that survival of sperm shows biphasic curve where at a very slow velocity ice will grow in a planar form which will kill all cells. At higher velocity ice crystals will form secondary branches and survival will increase, however at higher velocity (i.e. 300 μ m/sec) ice will start to form "needle-like" ice crystals which will decrease PTM, but in a higher velocity will permit very high survival (fig. 2) depending on the space between the ice crystals (4). Finally, at very high velocity (i.e. >3000 μ m/sec), directional solidification will not occur and survival will decrease.

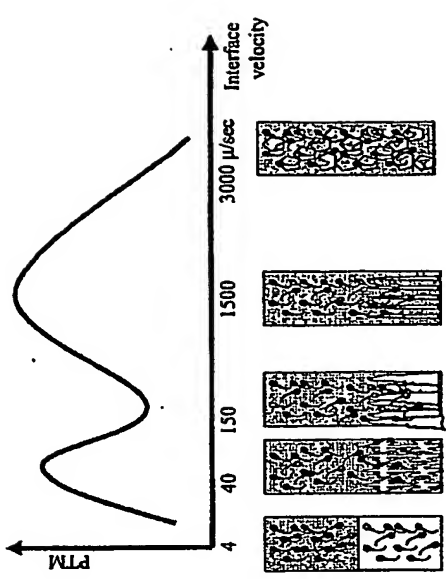


Fig. 2 Effect of interface velocity (V) on ice crystals morphology and sperm post thaw motility (PTM)

Heat transfer problems associated with large volume freezing

In a conventional slow-freezing method, temperature of the chamber is dropped in a controlled stepwise manner. This method is based on using multidirectional (equiaxed) heat transfer to achieve a rate of temperature change in the sample that depends on the thermal conductivity and geometrical shape of the container and of the biological material within it (5). The thermal gradient within the sample is determined implicitly by the temperature of the chamber and the thermal conductivity of the materials of the sample, and is not directly controllable. Furthermore, the ambient temperature gradients within the freezing chamber and the unreliability of temperature recording measurements (6) add to the difficulty of achieving the optimal cooling rate in a large volume sample.

Cryobanking of large volume semen

Each of the ejaculates was tested for semen concentration and motility (>70%) before dilution. We used AndroMed[®] (minitub, Flügplatzstrasse, Germany) for the semen dilution to have a final concentration of 1.5×10^6 sperm/ml. Freezing of a whole ejaculate was done in a special test tube (12ml) in which the central part is a hollow channel. Heat transfer is opposite to the test tube movement and is parallel to the tube length axis (fig. 1). The empty channel in the middle of the large test tube facilitate directional freezing and rapid thawing in the inner side of the test tube. Sperm PTM after freezing in a large volume was very high. We found a survival rate of 90-100% (normalized PTM) in the two bulls we cryopreserved in the MTG technique. These results were superior to MTG freezing using mini straws (data not shown), which suggest the benefit of using MTG freezing of large volume for sperm cryopreservation. Results shows a 75±5% post thaw motility after freezing a 12ml test tube and 50±5% after second freezing/thawing in a mini-straws, respectively. Controlled vapour freezing showed a 60±10% post thaw motility which were lower than the results after MTG freezing of mini straws.

The large volume freezing may be very useful for cryobanking of bull semen, for example, AI centre that have a bank of 10,000 straws which are made from 25 ejaculates (400 straws /ejaculate). We calculated that these 10,000 straws will fit into 13 goblets (750 straw/goblet). In comparison, when we freeze a large volume (12ml test tube) the 25 ejaculates will be frozen in 25 test tubes which will be stored only in 2 goblets. This means that we need 6.5 time more goblets using straws in comparison to test tube freezing. In this case, the present method gives a capability to have a bank of "waiting bulls" in some of the AI centers which presently do not use a semen cryobanking. In addition, this method will save money in labour and consumables (filling, printing, LN for freezing and for storage etc.).

In conclusion, the MTG technique could be very useful for large volume cryopreservation and double freezing for sperm cryobanking.

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multigradient tubes

Multigradient

Notes from interview with Amir Arav, 26 June 27-06-2002

Based on US 5,873,254.
Also based on the cryopreservation part of US 60/345,643.

Equine: most stallion sperm can't be frozen. We get higher recovery than others.

Largest volume of semen frozen in prior art in test tube = 5 ml. (larger volume of pig semen has been frozen in bags.)

Our innovation: large volume freezing.

Horse prior art: 0.5 ml. One insemination needs four to eight 0.5 ml semen samples.

Why large volume?
For example, for breeding cattle. Genetics breeding. Test young bulls for production. Collect semen. Inseminate heifers. Heifers give birth to heifers. Inseminate the daughters. See how much milk the daughters give. This takes 4 to 5 years. Only one out of every 14 candidate bulls is selected and something may happen to the best bull during the 4-5 years, like the bull may die. Therefore, need to put semen in bank. Store 10,000 to 50,000 0.25 ml straws per bull. It takes 25 days to collect 10,000 straws, and lots of liquid nitrogen for storage.

For volume of one ejaculate: We do one test tube, 12 ml sample, 1:2 dilution instead of 400 to 600 straws, 1:10 dilution. At the end of the 4-5 years, the selected bull's semen is thawed and refrozen in regular straws.

Prior art concept: can't freeze samples bigger than 0.5 ml, can't freeze concentrated sperm.

Prior art: 50,000,000 sperm cells per ml.
Us: 500,000,000 sperm cells per ml.

Equine and boar semen freezing without centrifugation.

Prior art: do centrifugation of semen to remove seminal plasma before freezing. Sperm concentration in semen is low in stallion and boar: 50,000,000 to 600,000,000 sperm cells per ml.
To remove plasma: centrifuge and wash the sperm. This damages the sperm. Then add extender to get concentration needed for insemination: 1,000,000,000 to 6,000,000,000 sperm cells total.

Us: dilute the plasma. Go down to 20,000,000 sperm cells per ml. Then need 50 ml for one insemination. Large volume freezing allows freezing one insemination (10 ml x 5 or 50 ml x 1) at once.

How?
Multigradient freezing of rotating test tube. Rotate the tube around its longitudinal axis during the freezing.

multi-grooved nozzles

Advantage: mix the solution in front of the ice front. This dilutes the concentration of salt being expelled from the ice.

Rotating tube also used for warming.

Rotating tube also used for freezing partly filled test tube. Spread solution in annulus to get high surface area.

Rotating the tube gives better thermal contact between the solution and the metal heat exchanger through the walls of the test tube.

The thermal contact with the block is always best on the bottom side of the tube, and that thermal contact gets spread around when the tube is rotated.

If the tube is partly filled with a sample, you wind up with frozen sample on the wall of the tube and air along the axis of the tube. That's better if then you want to lyophilize (freeze dry) the sample.

Rotating the tube keeps the sample mixed and homogeneous during freezing.

Alternative: hollow (double walled) test tube.

Piston sits inside central channel of test tube and removes heat from the central channel. More efficient heat exchange during cooling.

For heating: put hollow test tube in water bath so circulating water flows through the central channel. Warming rate in center of tube and on outside of tube is the same. Unlike US 3,871,254 seeding, need to inject liquid nitrogen at bottom of test tube. A section at the base of the test tube is arranged to exclude sperm but include liquid, for example by putting glass balls as liquid trap in base of test tube, so only the liquid is frozen for seeding.

Hollow tube can be glass or plastic. Hollow tube need not rotate.

Hollow tube also has a roughened section on side for manual marking.

Warning on thawing small straws (0.25 cc or 0.5 cc samples). Faster is faster, but can't go fast from liquid nitrogen temperatures or heat stress will crack the sample.

OTOH, overheating leads to denaturation, and warm cryoprotectant can damage sperm.

Faster is better to prevent recrystallization at -10°C. Machine has one block, at uniform high temperature: 38°C to 100°C, with 90°C being preferred. Put straw in hole through block. Move straw through the block via the hole at constant velocity. 6mm/sec optimal. Block is 2 cm thick, so 3.3333 second duration. Then out to ambient air. From when straw leaves liquid nitrogen to when straw emerges from block should be less than 50 seconds, preferably about 30 seconds.

Pull straw from liquid nitrogen: goes up to -30°C just by being in ambient air. Put straw through block: in 3 seconds, go to room temperature.

Another way to warm the test tube: like the straws.

Another way to warm a prior art test tube:

multigradient noise

Plunge into a water bath warmer than 37°C, with 70°C preferred. So no contact between ice and walls. Then drop contents of test tube into high volume pre-warmed dilution (insemination) solution. Mix. Alternative: keep the test tube in hot water while "stirring" with the test tube to get uniform thawing.

Freezing and thawing test tube with controlled rolling (rotating) system

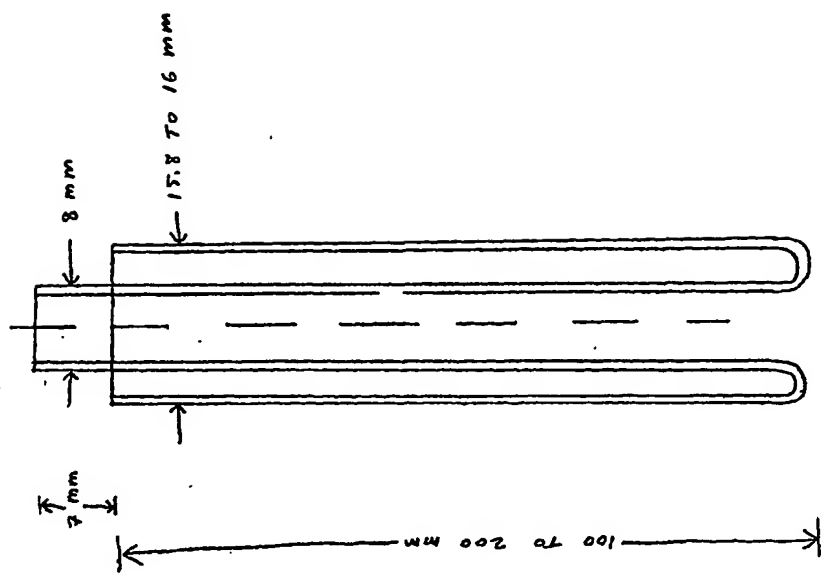
Inventors: Arav Amin, Meir Uri

We developed a device which has controlled rolling system of round container (i.e. test tube) during freezing and thawing.

The advantages of this device are:

1. A better heat transfer between the container and the copper blocks.
2. A formation of air bubble in the centre of the contained a thin layer on the wall of the container.
3. A controlled propagation of ice crystals in parallel to the container wall.
4. A continuous mixing of the solution during the freezing and thawing.
5. A preparing of large surface for the purpose of sublimation for freeze drying.

CROSS-SECTION OF HOLLOW TEST TUBE



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OBST BEACHIN STUD PAGE

DOUBLE FREEZING SECOND EXPERIMENT.

After the first experiment we had the best results for freezing 12 ml. tube was 12 second electrode. In the second step we to check a big number of bulls based on the results of the first experiment for better comparison in the experiment we chose 0.25 ml. tubes with the same sperm concentration 20K sperm for all treatments.

MORTALITY %

	C. Ryan	L William	H. Conrado	Alejo	Luzan	W. German	C. Dalton	A
First freezing	70	60*	60	60	60	60	60	71
After first freezing before	60	60	60	60	60	60	60	6
After first freezing D. Coro (control)	65	65	60	70	65	70	65	75
After second freezing straw 1ml	25	50	15	50	50	40	50	4
After second freezing straw 0.25 ml	10	45	40	40	40	50	40	3

Conclusion as the average of the results show:

- 1) After first freezing D. Coro machine is better than first freezing in 12 ml. tubes in 12% machine 70% vs 61.1%.
- 2) After second freezing D. Coro machine shows a 20% better results the D. Coro machine 40.0% vs 30.0%.
- 3) There are individual differences between the bulls in the freezing capacity of the semen and between the two freezing techniques, for example the bull Conrado frozen better in D. Coro machine 47% vs 15%.
- 4) We can conclude that it is possible to perform a field trial in the second freezing with both freezing techniques, with these bulls that I carried out with variable from 40% to 50% in D. Coro and 40% machine in the second freezing the best bulls for these techniques was are: L. William, Alejo, L. Luzan, W. German and C. Dalton.

ADDED BEACH MOTILITY, DEADLINE RATIO AND MEMBRANE INTEGRITY AFTER THAWING

The experiment:

The aim of this experiment was to test the best thawing time for 0.5 ml aliquots of 90 degree Cebuffer. For aliquots of 0.25 ml were found to be 2 seconds in 60 degrees celscius for best thawing time. At the beginning of each experiment were tested two, three of four strains of 0.5 ml and thawed at different thawing time. Two rates were applied for the assessment of the strains:

- 1) Viable motility after rate of incubation at 37C
- 2) Propidium iodide DAPI test for dead/live ratio dead cells/red colour and live cells blue colour.

The results:

Motility of strains at different thawing time of 0.5 ml aliquots at 1600

data	rate	thaw	3	4	5	6
24.04.02		Madonabean				
30.04.02		Shaker	60	60	60	30
21.04.02		Lucky	60	60	60	60
01.04.02		Shaker	35	45	60	60
02.04.02		Broker	60	40	75	60
03.04.02		Principal	20	30	50	60
07.04.02		Lucky	60	70	50	50
10.04.02		Courier	60	70	50	50
17.04.02		Shaker	60	60	40	50
18.04.02		Courier	30	60	50	50
Avg			41.3	47.2	60.0	47.1

Propidium iodide test (dead/live) the strain ratio

data	rate	Madonabean	Shaker	Lucky	Broker	Principal	Courier	Shaker	Courier	Avg
24.04.02										
30.04.02			61	65	65	67				
31.04.02			55	54	55					
01.04.02			63	73	73					
02.04.02			61	70	60	71				
03.04.02			51	68	77	75				
07.04.02			55	61	60					
10.04.02			65	64	65					
17.04.02			77	63	62					
18.04.02			60	72	63					
Avg			61.8	61.3	70.8	69.3				

Results: The average of the results showed a better motility and propidium iodide test 60% and 70.4% respectively after 6 sec. heating to water temperature of 60C for 0.5 ml strains. Low results were find in 3, 4 and 6 sec. thawing. Conclusion: The best results for both test were the sec. thawing at 90C, for 0.5 ml strains.

SOLVENT BREATHABILITY, DEGRADATION RATIO AND MEMBRANE INTEGRITY AFTER FREEZING

The second experiments based on the first experiments of the solvent breath that normally produce solvent for testing was tested in two temperatures 27°C and 0°C and two volumes of solvent 0.25 ml and 0.5 ml. In order to find the best solvent for freezing and thawing solvent in the ATC020 and ATC020-450 machines. The heating time of 10 sec at temperature of 37°C and the time to 0.25 ml solvent to 0.5 ml solvent and the freezing time for temperature of 0°C was 2 sec to 0.25 ml solvent and 5 sec to 0.5 ml solvent.

The freezing velocity was 0.5 ml/min and 1.5 ml/min. With a heating time of 0.5 sec and for the 0.25 ml solvent the freezing velocity was 2.0 mm/sec. With a heating time of 1.5 sec and for the 0.25 ml solvent the heating velocity was 1.0 mm/sec. With a heating time of 1.5 sec.

1) We used methyl acrylate for the estimation of the solvent stability.
 2) Propylene oxide was for solubility test
 3) 0.1% Tari for membrane integrity, the solvent included in a complete solution for 40 min when it is related with a fluorescent dye that can identified the damaged points, we used propylene oxide again.

The results:

date	bill	Methyl acrylate			Propylene oxide		
		0.25 ml. 37°C/min	0.25 ml. 0°C/min	0.5 ml. 37°C/min	0.5 ml. 0°C/min	0.25 ml. 37°C/min	0.25 ml. 0°C/min
23/6/02	Darzi M	60	60	60	60	60	60
24/6/02	Meddourneem	50	50	50	50	50	50
24/6/02	N. Omer	60	60	60	60	60	60
30/6/02	SHAW	60	60	60	60	60	60
31/6/02	Lurcy	60	60	70	60	60	60
01/7/02	Shaker	60	60	60	60	60	60
02/7/02	Bosher	60	60	60	60	60	60
02/7/02	Prashaw	60	60	60	70	70	70
07/7/02	Lurdy	60	60	60	60	70	70
10/7/02	Coures	60	70	70	70	70	70
17/7/02	Bosher	60	70	70	70	70	70
18/7/02	Coures	60	60	60	60	60	60
AVG		67.4	64.0	67.8	64.3		

date	bill	Methyl acrylate			Propylene oxide		
		0.25 ml. 37°C/min	0.25 ml. 0°C/min	0.5 ml. 37°C/min	0.5 ml. 0°C/min	0.25 ml. 37°C/min	0.25 ml. 0°C/min
23/6/02	Darzi M	60	60	60	60	60	60
24/6/02	Meddourneem	62	60	60	60	60	60
24/6/02	N. Omer	45	40	70	70	70	70
30/6/02	Shaker	60	60	60	60	60	60
31/6/02	Lurdy	60	60	60	60	60	60
01/7/02	Bosher	60	60	60	60	60	60
02/7/02	Prashaw	60	60	60	60	60	60
07/7/02	Lurdy	70	70	70	70	70	70
10/7/02	Coures	70	70	70	70	70	70
17/7/02	Shaker	77	77	77	77	77	77
18/7/02	Coures	61	60	60	60	60	60
AVG		62.3	64.4	62.7	64.3		

ORT 40 ml. incubation (paraffin integrity test)						
		0.20 ml.	0.20 ml.	0.05 ml.	0.05 ml.	
		370.1mln	300.2mln	370.1mln	300.2mln	
21.05.02	Darbie	45	37	55	49	
24.05.02	Kaufmann	51	35	45	43	
28.05.02	N. Orlay	59	35	45	37	
30.05.02	Shuler	40	42	51	57	
31.05.02	Lucky	67	44	32	35	
01.06.02	Shuler	60	67	44	39	
02.06.02	Boche	45	51	35	65	
03.06.02	Priest	52	35	55	48	
07.06.02	Lucky	35	35	52	47	
10.06.02	Coulier	69	51	34	61	
17.06.02	Shuler	70	53	52	70	
18.06.02	Coulier	53	44	25	41	
	Avg	49.8	48.0	38.6	48.3	

The results: The best results in mobility were for the high (incubation temperature (30°C) for the volume 0.20 ml) and 0.5 ml (65% and 65.5%) and the lower results were for the low temperatures (37°C) for 0.20 ml and 0.5 ml with 37.5% and 67.4% respectively.

For the Paraffin Incubation test (dead/alive ratio) the best results were for 0.20 ml and 0.5 ml and the lowest thermal stress obtained at lower temperatures 370, 0.2 and 0.5 ml respectively.

For the ORT test (membrane integrity of the plasma cells) the best results were for 0.20 ml and 0.5 ml extremes tested at high temperatures 30°C, 45.0% and 45.2% respectively and lower results were found for 0.20 ml and 0.5 ml extremes tested at low temperatures 37°C, 45.5% and 35.7% respectively.

Conclusion

The lower thermal at high temperatures showed an improvement in mobility, Paraffin test, ORT test and ORT test (membrane test) due to the damage caused to the plasma cells when the thawing is done at high temperatures (30°C) (not thawing). The volume of 0.5 ml always show a slightly better results in all three test than for 0.20 ml always when thawed at high temperature (30°C). The results above on advantage for big volumes. This seems to give better protection to the plasma cells under drastic changes in temperature, no differences were found for 0.20 ml and 0.5 ml always when thawed at low temperature (37°C) except for the ORT test, the better results were for 0.20 ml always than 0.5 ml always 45.5% vs 35.7% respectively.

Flow function comparison between 0.5 ml/min. Straws frozen in MTC-450 vs. 0.25 ml/min. straws frozen in MTC-450 and Digit Cool machine as controls.

The aim of this experiment was to compare 0.5 ml/min. straws frozen in MTC-450 in three different velocities (1200, 1500, and 1800 mm/sec vs. 0.25 ml/min. at 2000 mm/sec). The best results for the machines. The temperatures for both machines were 5°C for the first block and 30°C for the second block, the freezing temperatures of the straws was 37°C.

The data is from eleven blocks, that were frozen for internal freezing. The concentration of the serum was 20X10% per straw, the serum was diluted at room temperature and each drop of serum at the temperature of 6°C, the straws were filled and then frozen in the different machines.

Results

The mobility of要素, frozen in MTC-450 at different velocities compared to MTC-450 and Digit Cool machines

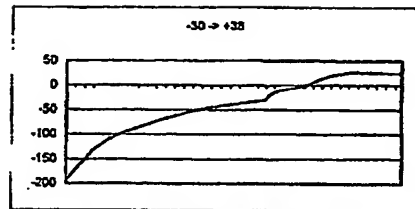
Fluid	Rate element	V-1200		V-1500		V-1800		V-2000		Digit Cool	
		MTC-450	MTC-450								
Hb solution	60	30	50	5	5	50	50	50	50	50	50
Draemer	80	60	75	5	5	75	65	65	65	65	65
Colura	80	40	65	5	5	60	40	40	40	40	40
Protein	60	50	75	2	2	75	60	60	60	60	60
Loeffler	70	60	60	5	5	60	40	40	40	40	40
Pfele	80	65	70	55	55	70	60	60	60	60	60
Cordian	70	50	60	5	5	65	50	50	50	50	50
C. Chedier	80	50	45	5	5	50	50	50	50	50	50
Pasha	40	30	30	2	2	50	50	50	50	50	50
C. J. Hirsch	50	60	60	2	2	75	60	60	60	60	60
S. Nation	60	60	60	10	10	40	50	50	50	50	50
AVG	72.7	46.0	65.4	9.2	9.2	60.0	48.8	48.8	48.8	48.8	48.8

Conclusions

The best results were found for the MTC-450 at a velocity of 2000 mm/sec. 60% and for MTC-450 at a velocity of 1500 mm/sec. with 50% mortality, better results were found for Digit Cool machine conventional freezing with 40.5% and for MTC-450 at a velocity of 1200 mm/sec. with 45% mortality, the worst results were obtained for MTC-450 at a velocity of 1000 mm/sec. with 9.2% mortality.

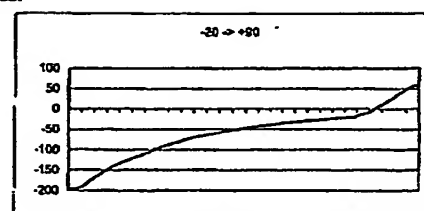
Bull sperm thawing in Drawing machine

#	T-#	Vel/time	Air time	Microscope							
				% motil	SYBR	PI	SYBR	PI	SYBR	PI	
1	80	3 sec.		10	70						
2	80			15	70						
3	80			20	50						
4	80			25	60						
5	90			25	55	89	53	56	77	70	112
5	90			30	70						
7	80			30	70	30	15	28	15	25	14
6	80			30	70	55	40	41	30	25	13
9	80			35	70	35	25	40	25	43	25
10	80			40	45						
11	80			50	15						

Bull semen thawing ...**-30 \rightarrow +38**

#	Algorithm	% motility
1	-198 > -24(20s) > 38	40
2	-198 > -24(20s) > 38	65
3	-198 > -24(20s) > 38	70
4	-198 > -24(20s) > 38	75
5	-198 > -24(40s) > 38	60
6	-198 > -24(40s) > 38	40
7	-198 > -24(40s) > 38	65
8	-198 > -24(40s) > 38	70

Average: 62.14286

1. Using water bath**-20 \rightarrow +38**

#	Algorithm	% motility
1	-198 > -24(20s) > +80(2s) > 38	70
2	-198 > -24(20s) > +80(2s) > 38	75
3	-198 > -24(20s) > +80(2s) > 38	75
4	-198 > -24(20s) > +80(2s) > 38	75
5	-198 > -24(40s) > +80(2s) > 38	75
6	-198 > -24(40s) > +80(2s) > 38	75
7	-198 > -24(40s) > +80(2s) > 38	75

Average: 74.28571

2. Using new thawing machine

#	Air time	Hot block Temp.	Time (sec)	% motility
1	10	60	3	70
2	15	60	3	70
3	20	60	3	60
4	25	60	3	60
5	25	60	3	55
6	30	60	3	70
7	30	60	3	70
8	30	60	3	70
9	35	60	3	70

Stallion semen results

11.06.02

Stallion	Pre-freeze motility	Chilled semen after 30hrs				Planer Straw				MTG Tube			
		%motility	AO/PI %alive	ORT	%motility	AO/PI %alive	ORT	%motility	AO/PI %alive	ORT	%motility	AO/PI %alive	ORT
Nomrod	65	60	58.8	49.2	50	38.5	25.3	55	46.8	30.0			
Jet set	80	70	59.4	47.1	65	54.0	41.6	65	76.5	50.5			
William Curtis	50	60	62.3	38.0	20	25.4	18.1	40	55.0	37.7			
Libra K	60	40	46.4	24.6	25	28.3	12.2	60	59.0	34.3			
Samhba	80	70	70.0	59.1	60	51.4	33.0	70	68.0	40.3			
Mean	67.0	58.0	57.3	43.2	42.0	39.5	26.0	66.0	61.1	38.6			

27/06/02 15:31 27/06/02 15:31

Dr. N. Friedman *** 0351575-5 090

Station search results

10.06.02

Station	Pre-freeze motility	Chilled semen after 30hrs				Planer Straw				MTG Tube			
		%motility	AOI/PI %live	ORT	%motility	AOI/PI %live	ORT	%motility	AOI/PI %live	ORT	%motility	AOI/PI %live	ORT
Jet Set	70	40	48.4	24.8	60	49.3	27.1	60	58.1	46.2	60	58.1	46.2
Cheron	70	60	48.1	46.2	40	38.2	21.0	60	41.0	26.9	60	41.0	26.9
Rubek	80	80	69.2	57.9	60	54.0	24.0	60	51.4	32.1	60	51.4	32.1
Nemrod	60	60	64.1	52.1	30	34.3	12.0	60	49.7	21.5	60	49.7	21.5
Mean	70.0	60.0	66.7	45.3	45.0	44.2	23.5	55.0	60.1	31.7			

Dear Udi

I hope this information is not too late, I have just received your e-mail.
Please find attached the results for the latest MTG protocol for stallions. I am applying three tests for post thaw evaluation, namely Osmotic Resistance Testing (a membrane strength stress test), Acridine Orange/Propidium Iodide (a membrane viability test) and motility. None of these tests are unique to us.
The freezing extender I am using contains the following:

Glucose monohydrate 15g
Trisodium citrate 0.925g
EDTA 0.925g
Sodium hydrogen carbonate 0.3g
Lactose 55g
Lauryl sulphate 0.375g
Lignospectin 1.0g
Gentamycin 1.25ml
Clarified egg yolk (centrifuged at 10000XG to remove fat) 200ml
Glycerol 30ml (3%)
Water to 1000ml

The clarified egg and low glycerol concentration makes this extender unique to us and could be regarded as specific or the MTG.

The freezing protocol I use is:

Manual sealing
5°C start temp
-50°C end temp
1.0mm/second velocity.

A range of velocities can be applied from 0.6mm/second-3.0mm/second. These should all be protected. Also the start temp can be altered to a range between 25°C-5°C and the end temp can be between -5°C-100°C.

I think you are familiar with all other aspects (tubes etc.). Please do not hesitate to ring me if you need any other assistance. I am here for most of the day.

Regards
Matt

The following section of this message contains a file attachment prepared for transmission using the Internet MIM/E message format. If you are using Pegasus Mail, or any other MIM/E-compliant system, you should be able to save it or view it from within your mailer. If you cannot, please ask your system administrator for assistance.

8.06.2002

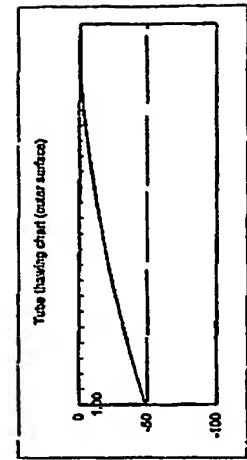
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DR. M. PRIDMAN *** B.I.U. 601391575 - 06

Stallion semen results

Stallion	Pre-freeze motility	Chilled semen after 30hrs				Plano Straw				MTG Tube			
		%motility	AO/PI	%live	ORT	%motility	AO/PI	%live	ORT	%motility	AO/PI	%live	ORT
Memphis	60	50	47.9			60	63	29.6	60	57.4		37.1	
Oberon	50	50	54.2			10	7.2	0	10	6.1		0	
Nemrod	55	50	56.1			25	29.5	20.0	40	41.5		29.3	
Rubek 1	60	50	45.7			30	29.1	13.4	50	54.8		35.4	
Rubek 2	60	50	53.8			30	34.6	16.4	50	56.9		41.1	
Eagle	70	60	68.4			25	18.1	12.1	60	52.6		49.8	
Lagos	60	60	69.1			55	57.6	29.8	60	63.4		50.0	
William	60	55	65.0			45	52.3	39.4	60	58.7		39.8	
Mean	69.4	63.1	68.3			33.8	35.2	20.1	48.8	49.2		35.3	

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DR. M. FRIEDMAN *** BILL
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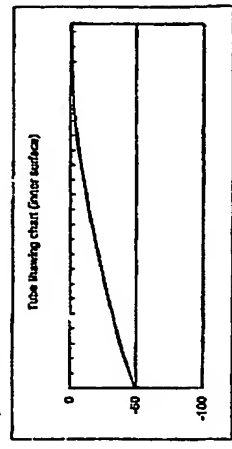
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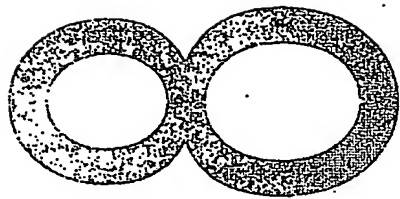
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DR. M. FRIEDMAN 60394575 800

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Oath/Declaration, Small Entity
and Power of Attorney



Level - 2
Version 1.1
Updated - 8/01/01

Int'l

ASSIGNMENT

For good and valuable consideration, the receipt and sufficiency of which is hereby acknowledged, a
undersigned:

Anil Arav

Mail Us

hereinafter called the "assignor(s)", hereby sell(s), assigns(s) and transfer(s) to:

Interface Multigrid Technology Ltd.
31 Hamazem St.
Ness Ziona 76400
Israel

(hereinafter called the "assignee(s)"), his/her successors, assigns, nominees or other legal representatives, all
Assignor's entire right, title and interest in and to the invention entitled:

EMBRYO GUARD

described and claimed in the following patent applications:

U.S. Provisional Application identified as Attorney docket No. 79/14 and executed the same date as
assignment;

and in and to said Patent Applications, and all original and related Patents granted therefor, and all division
continuations thereof, including the right to apply and obtain Patents in all other countries, the priority rights
under International Conventions, and the Letters Patent which may be granted thereon;

Signed and sealed this 22 day of June 2002


Anil Arav
14/06/2002

Mail Us
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18.05.02-19.05.02

Results

Stallion	Pre-freeze motility	Chilled semen after 30hrs				Planer Straw				MTG Tube				Status
		%motility	AO/PI %live	ORT	%motility	AO/PI %live	ORT	%motility	AO/PI %live	ORT				
C. R. Gold	25	0	4	0	5	1	0	15	21.6	3.4				fail-fail
Libra-K	80	20	19.4	12.1	20	40.0	33.3	35	43.7	29.4				fail-pass
Samhre	60	60	72.4	45.2	3	31.9	23.3	60	46.6	46.0				fail-pass
Libra-K	80	60	79.1	48.4	20	42.6	33.9	40	57.0	46.0				fail-pass
Mill Law	40	30	42.3	39.2	10	34.5	26.2	30	42.2	35.6				fail-pass
Jester	80	60	71	59.1	30	29.3	20.1	40	40.3	23.2				fail-pass
Rob Roy	80	80	80.2	78.9	60	75.2	66.3	80	76.1	64.1				fail-pass
Pall Mall	70	20	18.4	0	25	32	16.3	35	38.4	26.2				fail-pass
Jester	60	60	48.6	41.2	30	34.2	22.6	60	53	43.4				fail-pass
Drambro	70	50	55.2	40	20	24.3	11	40	49.4	41.5				fail-pass
Rubek	60	50	58.8	47.1	35	39.4	21.2	50	44.4	43.9				pass-pass
Rubek	60	60	52.3	51.1	50	49.8	35.9	60	49.2	40.2				pass-pass
Secundus	70	25	39.1	30.2	35	23.5	22	60	53.9	42.6				pass-pass
Schiller	80	60	71	49.8	60.0	43.5	34.7	60.0	64.6	51.2				pass-pass
Ludwig	80	40	39.4	28.4	35	35	28.9	40	38.6	29.4				pass-pass
Schiller	80	60	49.4	37.4	40	37.5	27.5	45	48.3	27.5				pass-pass
Secundus	80	50	62.8	50.1	40	43.2	38.2	50	51.4	37.9				pass-pass
C. R. Gold	0													
Mean	65.8	43.1	48.6	38.7	29.9	34.3	25.0	43.3	45.6	35.1				
Mean (fail-pass)	71.1	47.8	55.3	40.7	25.4	38.2	27.0	45.8	50.1	33.6				
Mean (pass-pass)	74.3	49.3	53.0	42.0	42.1	38.8	29.5	60.7	49.6	38.0				
					Straw	7/17 pass		Tube	16/17 pass					

Stallion semen results

To date 18.06.02

Stallion	Pre-freeze motility	Chilled semen after 30hrs			Pipette Size	NGT Tube	ORT	
		%motility	AOPt Xlive	ORT				
Samshire	60	0	4	0	3	21.3	23.3	60
Charming Gold	60	50	54.2	0	10	34.8	26.2	16
Sili Law	50	50	64.2	40	10	7.2	0	42.2
Oberon	50	50	65.2	40	10	26.8	12.6	8.1
Sazzus	50	50	65.2	40	10	24.3	11	41.1
Danzels	50	50	65.2	40	10	40.0	33.3	43.7
Libra K	50	50	65.2	40	10	42.8	33.9	57.0
Libra K	50	50	65.2	40	10	21.4	15.1	55.0
Williams Curtis	50	50	62.3	35.0	10	33	20.5	31.7
Eagle	62	62	65.4	0	10	14.1	12.9	42.8
Eagle	62	62	65.4	0	10	24.1	19	49.8
Libra K	40	40	46.4	24.6	10	28.3	12.2	59.0
Libra K	40	40	46.4	24.6	10	28.7	29.7	49.1
Memphis	50	50	55.1	0	10	29.5	20.9	41.6
Memphis	50	50	55.1	0	10	32	19.3	35.4
Nemrod	50	50	55.1	0	10	28.4	27.8	44.9
Rob Roy	50	50	55.1	0	10	29.3	20.1	46.3
Perth Hall	50	50	55.1	0	10	34.3	13.5	53
Charming Gold	50	50	55.1	0	10	34.3	22.6	43.4
Jester	71	71	56.1	0	10	34.3	13.5	49.7
Jester	48.6	48.6	41.2	0	10	34.3	13.4	21.5
Nemrod	64.1	64.1	52.1	0	10	29.1	13.4	54.8
Robek	45.7	45.7	53.0	0	10	34.6	15.4	55.9
Robek	53.0	53.0	53.0	0	10	21.9	17.1	57.1
Russell	39.4	39.4	28.4	0	10	35	24.9	38.6
Ludwig	61.1	61.1	64	0	10	47.2	30.7	53.8
Rob Roy	71.0	71.0	62.7	0	10	41.4	32.2	51.1
Rob Roy	50	50	50	0	10	36.4	26.2	34.5
Robek	32.1	32.1	30.2	0	10	25.5	2.2	44.4
Decimus	50	50	50	0	10	44.0	20.5	63.9
Nemrod	48.1	48.1	45.2	0	10	39.3	21.0	50
Oberon	49.4	49.4	37.4	0	10	37.5	21.5	41.0
Schiller	64.2	64.2	60	0	10	49.4	37.9	52.3
Schiller	65	65	68.4	0	10	49.7	36	53.4
Secundus	62.8	62.8	60.1	0	10	43.2	30.2	51.4
William	65.0	65.0	60.1	0	10	52.3	33.4	57.6
Jet Set	45.4	45.4	24.0	0	10	49.3	37.1	53.7
Memphis	47.9	47.9	52.5	0	10	53	35.5	57.4
Nemrod	52.5	52.5	49.2	0	10	49.8	35.2	50.8
Robek	50	50	50	0	10	49.8	35.2	49.2
Samshire	70.0	70.0	59.1	0	10	51.4	33.2	65.0
Williams Curtis	62.1	62.1	57.0	0	10	49.7	25.1	58.4
Lagos	62.2	62.2	57.0	0	10	43	35.1	54.8
Robek	57.4	57.4	49.8	0	10	52.3	24.0	63.4
Schiller	57.4	57.4	49.8	0	10	54.6	24.0	60.0
Jet Set	53.2	53.2	47.1	0	10	42.5	34.7	61.4
Rob Roy	53.2	53.2	47.1	0	10	54.0	41.6	61.2
					8	73.3	68.3	75.8
								50.8
								54.1

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DR. M. FRIEDMAN *** BIR 6-1575-00

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